Abstract—Early detection of AD is of vital importance in the development of disease-modifying therapies. This necessitates the use of early pathological indicators of the disease such as amyloid abnormality to identify individuals at early disease stages where intervention is likely to be most effective. Recent evidence suggests that cerebrospinal fluid (CSF) amyloid $\beta_{42}$ ($A\beta_{42}$) level may indicate AD risk earlier compared to amyloid positron emission tomography (PET). However, the method of collecting CSF is invasive. Blood-based biomarkers indicative of CSF $A\beta_{42}$ status may remedy this limitation as blood collection is minimally invasive and inexpensive. In this study, we show that APOE4 genotype and blood markers comprising EOT3, APOC1, CGA, and $A\beta_{42}$ robustly predict CSF $A\beta_{42}$ with high classification performance (0.84 AUC, 0.82 sensitivity, 0.62 specificity, 0.81 PPV and 0.64 NPV) using machine learning approach. Due to the method employed in the biomarker search, the identified biomarker signature maintained high performance in more than a single machine learning algorithm, indicating potential to generalise well. A minimally invasive and cost-effective solution to detecting amyloid abnormality such as proposed in this study may be used as a first step in a multi-stage diagnostic workup to facilitate enrichment of clinical trials and population-based screening.

I. INTRODUCTION

Alzheimer’s disease (AD) is the most common neurodegenerative disease accounting for over 60% of all dementia cases [1]. It is characterized in part by the accumulation of amyloid-beta ($A\beta_{42}$) plaques in the brain – a condition known as amyloid pathology – that is present long before clinical symptoms (cognitive) are apparent [2, 3]. No cure or disease-modifying treatment for AD currently exists. There are ongoing efforts in clinical trials to combat this challenge. Current clinical trials target individuals at the earliest stages of AD, where intervention is thought to be most likely successful, following the high failure rates of previous trials [4]. Amyloid screening is used in these trials to identify individuals with amyloid pathology and may therefore be at the early stages of the disease before symptom onset. It may also be beneficial in the future for population-based screening [5, 6].

Current validated biomarkers of abnormal amyloid accumulation include $A\beta$ positron emission tomography (PET) and $A\beta_{42}$ measurement in cerebrospinal fluid (CSF) [7]. Use of these markers is internationally recommended [3, 8]. However, PET scans are expensive and available only at specialized centres while lumbar punctures required for CSF testing are invasive.

Notwithstanding the invasiveness of CSF collection, there is growing evidence that CSF $A\beta_{42}$ may be an earlier indicator of AD pathology compared to $A\beta_{42}$ PET [9-11] and thus may be a more suitable biomarker for disease detection at the earliest stages. To mitigate the limitation of invasiveness posed by CSF-based amyloid testing, there is strong interest in identifying blood-based biomarkers reflective of amyloid status as would CSF. Such biomarkers may be used as a reliable initial step in a multistage diagnostic procedure.

A few studies [12, 13] have demonstrated the potential of blood-based markers predictive of amyloid status as measured by CSF $A\beta_{42}$ with area under receiver operating curve (AUC) reaching 0.88 (in 46 samples) and 0.81 (in 358 samples), respectively. However, the novel method employed by [12] in measuring the blood-based markers remains to be established and the results from [13] are yet to be validated in independent cohorts.

In this study, we explore the utility of blood-based proteins to predict CSF $A\beta_{42}$ status using support vector machines with recursive feature elimination (SVM-RFE) that has shown effectiveness in similar research domains [14]. We also give particular consideration to the robustness of identified markers, to enhance the likelihood of reproducing results since reproducibility of results is one of the challenges in AD blood biomarker discovery domain [15].

II. METHODS

A. Study data preparation

Baseline data of 566 individuals from Alzheimer’s Disease Neuroimaging Initiative (ADNI) cohort were obtained (http://adni.loni.ucla.edu). The data comprised blood-based measurement of 190 proteins analyzed on a Rule-Based Medicine platform and 3 other proteins (including homocysteine, $A\beta_{40}$, and $A\beta_{42}$). The data also included apolipoprotein E $\varepsilon4$ (APOE4) genotype, demographic and diagnostic information as well as CSF $A\beta_{42}$ levels of the individuals measured on the Lumines Xmap platform. Forty-four (44) of the proteins were later excluded due to missingness, leaving 149 proteins. Finally, data from 358 individuals remained after 208 were dropped.

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TABLE I.
DEMOGRAPHIC CHARACTERISTICS OF STUDY SUBJECTS

<table>
<thead>
<tr>
<th>Clinical Diagnosis</th>
<th>CN</th>
<th>MCI</th>
<th>ADD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of participants (n)</td>
<td>58</td>
<td>198</td>
<td>102</td>
</tr>
<tr>
<td>Age (mean, (SD))</td>
<td>75.11(5.77)</td>
<td>74.37(7.49)</td>
<td>74.86(7.88)</td>
</tr>
<tr>
<td>Gender, female (n, (%))</td>
<td>28(48.28)</td>
<td>65(32.83)</td>
<td>43(42.16)</td>
</tr>
<tr>
<td>Years of education (mean, (SD))</td>
<td>15.67(2.78)</td>
<td>15.80(2.99)</td>
<td>15.16(3.30)</td>
</tr>
<tr>
<td>APOE4 carriers (n, (%))</td>
<td>5(8.62)</td>
<td>106(53.56)</td>
<td>71(69.61)</td>
</tr>
<tr>
<td>Low CSF Aβ42 status (n, (%))</td>
<td>1(1.72)</td>
<td>147(74.24)</td>
<td>93(91.18)</td>
</tr>
</tbody>
</table>

CN: Normal control; MCI: Mild cognitive impairment; ADD: Alzheimer’s dementia; n: Number of subjects; SD: Standard deviation.

The objective here was to identify potential blood biomarker signatures predictive of CSF Aβ42 status, from which a signature can be selected based on robustness and performance. The measure of robustness was intended to be transparent and simple to evaluate. The method used is based on the approach proposed by Abeel et al. [16] with some modification.

Similar to [16], SVM-RFE [14] combined with ensemble technique was used to select features for signatures formation, while Kuncheva index (KI) [17] was used to evaluate robustness of signatures. SVM-RFE combines the embedded feature selection capability of linear SVM with backward feature elimination strategy of RFE. Absolute values of the weights (coefficients) the linear SVM provides is the contribution of each feature to the SVM hyperplane and may be used as a means of ranking the importance of individual features. A feature with a larger weight is regarded as one of higher importance, and one with a lower weight is considered less important.

RFE implements a backward feature elimination procedure that iteratively removes the least important features in the training data samples. The algorithm starts out by fitting the training data with all the available features to a linear SVM, then ranks the features according to their weights and eliminates the least important one(s). The training data is subsequently refitted to the linear SVM but with only the retained features. This process is repeated until all features have been eliminated or a desired number of features to retain is attained. Finally, each feature in the training data is assigned an overall rank \( r \) (an integer with 1 as minimum and dimension of training data \( D \) as maximum) according to the observed feature contributions, with most significant features assigned lowest ranks.

SVM-RFE with ensemble learning is implemented to improve the robustness (stability) of feature subset selection by SVM-RFE. In this approach, \( k \) different subsamples of the original dataset (of \( D \) dimension) are generated using random sampling without replacement, each subsample containing only a slight variation (\( p \) samples) of the original dataset. For each subsample (in the \( k \) subsamples), \( b \) bootstrap samples are generated. SVM-RFE provides with a specified signature size \( s \) as a stopping criterion is then applied to each bootstrap. The rank of each feature in \( D \) as well as the AUC performance (AUC\(_{OO} \)) of the selected features on the out-of-bag samples is recorded. A candidate signature of size \( s \) is subsequently selected according to an ensemble ranking \( R \) obtained by aggregating \( r \) over all \( b \) bootstrap samples as shown in (1).

An estimate of the generalization performance of the signature is obtained by training the linear SVM on the subsample and its performance evaluated on the \( I-p \) held out samples. Ensemble method of generating signatures has shown to improve robustness and classification performance compared to simply applying SVM-RFE directly to subsamples [16]. In addition to the approach proposed in [16], we carried out a repeated stratified cross-validation of the candidate signature on the corresponding subsample as a supportive evaluation of the signature’s classification performance.

\[
R = \sum_{i=1}^{b} w_ir_i
\]  \hspace{1cm} (1)

The weight \( w_i \) is bootstrap-dependent. It takes either of two values depending on the chosen aggregation method. In the complete linear aggregation (CLA) method, \( w_i \) is set to 1, while \( w_i = 1 - \text{AUC}_{OO} \) in the complete weighted aggregation (CWA) strategy. The two methods were explored in this study albeit CWA was shown to be marginally better than CLA in [16].

To evaluate the robustness of the \( k \) candidate signatures, a stability measure defined by the Kuncheva index (KI) [17] shown in (2) was applied.

\[
\text{KI} = \frac{m - (s^2/k)}{s - (s^2/k)}
\]  \hspace{1cm} (2)

KI with range \([-1, 1]\) measures the similarity between two signatures. \( m \) is the number of features common to both signatures. The greater the value of KI, the larger the number of common features. A negative index indicates that feature intersection is mostly due to chance. The overall stability \( \text{KI}_{\text{tot}} \) of a signature can be defined as the average of all pairwise similarity comparison between the signature and rest of the \( k-1 \) signatures.

\[
\text{KI}_{\text{tot}} = \frac{\sum_{i=1}^{k-1} \text{KI}_i}{(k-1)}
\]  \hspace{1cm} (3)

C. Implementation

The robust biomarker selection task was implemented in python programming language. The machine learning subtasks were conducted with the scikit-learn package. Codes are available at https://github.com/chimastan/robust-blood-based-signature-of-csf-abeta42-status. The values of \( k, b, \) and \( p \) used were 500, 50, and 0.8, respectively, considering the recommendations by [16]. Cross-validation fold used was 10-fold with 10 repetitions with samples stratified according to...
the target label distribution. The C parameter for the linear
SVM was set to default (C=1). In the RFE, number of features
to eliminate per run was set to 20% of the total available
features to improve speed of processing.

III. RESULTS

A. Potential robust signatures

We realized several potential signatures with different
levels of classification and stability performance for
prediction of CSF Aβ42 status. Fig. 1 illustrates the variation
between signature size s and the average cross-validated AUC
as well as average Kltot over the 500 subsamples. It can be
seen that the average AUC gradually increased with
increasing s up to a point (s=8) and then declined, while
stability steadily dropped with increasing s. The results of
CWA and CLA ensemble methods were largely equivalent as
shown in Fig 1. Thus all further reports are based on results
of the simpler CLA method. Consideration of potential
signatures was also limited to ones consisting of 5 biomarkers, being that stability remained moderate at s=5 while the increase in average AUC beyond that point was minimal.

A total of 229 unique candidate signatures were obtained
from the 500 subsamples. We then identified the top 10
signatures with best values of stability Kltot (ranging between
0.67 and 0.61) and subsequently carried out further analysis
to aid making a final selection.

B. Final selection of signature

We conducted additional analyses with similar approach
as in II(B) but with s limited to 5 and random forests (RF)
used as the machine learning algorithm. Therefore in this
case, RF-RFE was applied instead of SVM-RFE. The number
of trees per forest was set to 2000, each forest containing a
maximum of D^{3/4} features as recommended in [18]. The
purpose was to obtain candidate signatures with best Kltot
values and compare them to the top 10 realized earlier with
SVM-RFE. This would allow identifying signatures whose
classification and stability performance may be agnostic to
type of machine learning algorithm and thus likely to
generalize better. With the RF-RFE, we realized 169 unique
potential signatures and identified the top 10 with best
stability values. A comparison of the signatures with ones
obtained with SVM-RFE implicated one signature as
common. The signature consists of APOE4 genotype,
etoxin-3 (EOT3), apolipoprotein-C1 (APOC1) and
cromogranin-A (CGA), and Aβ42. The signature achieved
0.64 stability (Kltot) value. Average AUC, sensitivity,
specificity, negative predictive value (PPV) and negative
predictive value (NPV) for the repeated 10-fold cross-
validation were 0.85, 0.84, 0.63, 0.83 and 0.67, respectively.
The average values on the unseen held-out samples were 0.84
AUC, 0.82 sensitivity, 0.62 specificity, 0.81 PPV, and 0.64
NPV, respectively. Contribution of individual biomarkers to
the classification performance of the signature is as shown in
Fig. 2 with APOE4 unsurprisingly making the most
contribution.

IV. DISCUSSION

In this study, we investigated the utility of blood-based
signature predictive of CSF Aβ42 status with a robust
performance. We showed that APOE4 genotype and levels of
four proteins predicted CSF Aβ42 status with high AUC. This
is the first study to demonstrate a signature with a stable
performance beyond a single machine learning algorithm. It
is a positive indicator of the signature’s potential to
generalize.

Compared to existing studies, four out of the five
predictors (APOE4, CGA, Aβ42 and EOT3) in the signature
were implicated in a multi-marker panel from a recent study
[13] as predictive of CSF Aβ42 status with RF. A number of
studies have shown evidence of association between some of
the identified markers and AD. In line with our observed
prominent contribution of APOE4 in the identified signature,
it is the strongest and most prevalent genetic risk factor for
late-onset AD and considered as a possible therapeutic target
[19]. Serum and CSF but not plasma levels of EOT3 have
been shown to be dysregulated in individuals with AD [20].
APOC1 genes, in combination with APOE4, are suggested to
play an important risk factor role in AD [21, 22]. However,
association between plasma levels of APOC1 and AD has not
been evidenced. CGA has an amount of co-localisation with
brain amyloid plaques [23].
Figure 2. Contribution of individual marker to classification performance of the selected signature. The contribution was determined from the feature weights of linear SVM, normalized by the largest weight during training.

Notwithstanding, CSF and blood levels of CGA have not been reported to be correlated. Interestingly, plasma and CSF Aβ42 have shown to be correlated in individuals with AD [24, 25]. This study has several limitations. All analyses were conducted with the ADNI cohort with its peculiarity such as age and level of education of participants. Distribution of individuals with abnormal CSF Aβ42 levels across the clinical groups (CN, MCI, and ADD) was biased, with nearly all samples belonging to the MCI or ADD group. This might have impacted our analyses as the individuals are likely to have developed other confounding conditions.

V. CONCLUSION

Early detection of AD is crucial to the future success of disease modifying therapies which are thought to be most effective at the earliest disease stages. This necessitates the use of early pathological indicators of the disease such as amyloid abnormality. A minimally invasive and cost-effective solution to detecting amyloid abnormality such as Aβ42 status, may provide a promise for Alzheimer's disease patient classification and early intervention. We anticipate that the findings of this study can be applied to the clinical trials and population-based screening of the disease.

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